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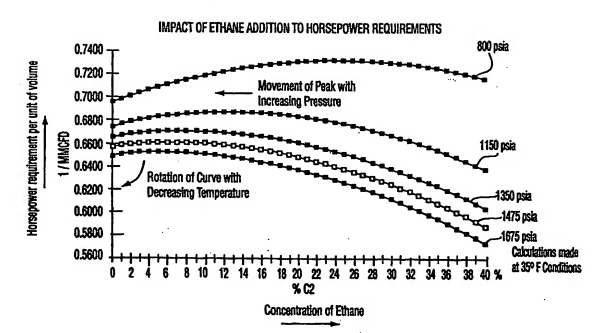
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(54) Title: PIPELINE TRANSMISSION METHOD



(57) Abstract

At pressures over 1,000 psia, it is advantageous to add to natural gas an additive which is a C_2 or C_3 hydrocarbon compound or a mixture of such compounds. Above a lower limit (which varies with the additive being added and the pressure), this results in a decrease in the amount of power needed to pump the mixture or to compress it.

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PIPELINE TRANSMISSION METHOD

Field of the Invention

This invention relates to the transfer by pipeline of mixtures which contain methane or natural gas.

5 Background of the Invention

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As is well known, methane is the largest component of natural gas, and usually accounts for at least 95% by volume of what is known as "transmission specification" natural gas. Other usual components are ethane (usually about 2%), propane (usually about 0.5%), butanes, pentanes and possibly hexanes (altogether amounting to less than about 0.3%), with the balance being nitrogen and carbon dioxide. In this disclosure, transmission specification natural gas will be hereinafter called "natural gas". For example, the natural gas as transmitted through the pipelines of TransCanada Pipeline Limited from Alberta, Canada to Ontario, Canada has typically the following percentage composition by volume:

15		Component (<u>Feed</u>
		Nitrogen Carbon Dioxide Methane Ethane	0.01270 0.00550 0.95400 0.01970
20	νĊ	Propane i-Butane n-Butane	0.00510 0.00170 0.00080
25		i-Pentane n-Pentane n-Hexane	0.00020 0.00010 0.00020

The relation between pressure, volume and temperature of a gas can be expressed by the Ideal Gas Law, which is stated as PV = nRT, where:

P = pressure of gas

V = volume of gas

n = number of moles of the gas

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R = the universal gas constant (which, as is known, varies somewhat depending on volume and temperature)

T = temperature of the gas.

If the equation is expressed in English units, the pressure is in pounds per square inchabsolute (psia), the volume is in cubic feet, and temperature is in degrees R (degrees Fahrenheit plus 460).

The Ideal Gas Equation does not give exactly correct results in actual practice, because gases are compressible. Gas molecules, when compressed, pack more tightly together than would be predicted by the Ideal Gas Equation, because of intermolecular forces and molecular shape. To correct for this, an added term, the compressibility factor z, can be added to the Ideal Gas Equation. This is a dimensionless factor which reflects the compressibility of the particular gas being measured, at the particular temperature and pressure conditions.

At atmospheric pressure or gage pressures of a few hundred pounds, the compressibility factor is sufficiently close to 1.0 so that it can be ignored for most gases, and so that the Ideal Gas Law can be used without the added term z. However, where pressures of more than a few hundred pounds exist, the z term can be different enough from 1.0 so that it must be included in order for the Ideal Gas Equation to give correct results.

According to the van der Waals theorem, the deviation of a natural gas from the Ideal Gas Law depends on how far the gas is from its critical temperature and critical pressure. Thus, the terms T_R and P_R (known as reduced temperature and reduced pressure respectively) have been defined, where

$$T_R = T$$
 T_c

$$P_{R} = P_{c}$$

where,

T = the temperature of the gas in degrees R

 T_c = the critical temperature of the gas in degrees R

P = the pressure of the gas in psia

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P_c = the critical pressure of the gas in psia

Critical pressures and critical temperatures for pure gases have been calculated, and are available in most handbooks. Where a mixture of gases of known composition is available, a pseudo critical temperature and pseudo critical pressure which apply to the mixture can be obtained by using the averages of the critical temperatures and critical pressures of the pure gases in the mixture, weighted according the percentage of each pure gas present.

Once a pseudo reduced temperature and pseudo reduced pressure are known, the compressibility factor z can be found by use of standard charts. One of these is "Compressibility Factors for Natural Gases" by M.D. Standing and D.L. Katz, published in the <u>Engineering Data Book</u>, Gas Processors Suppliers Association, 10th edition (Tulsa, Oklahoma, U.S.A.) 1987.

When the compressibility factor z of methane is read from the charts, it is found that the factor z is always less than 1.0 in normal temperature ranges (i.e. between about -40°F and 120°F) and that it decreases as the pressure rises or the temperature falls. Therefore, less energy need be used to pump a given volume of methane (measured at standard volume) at any given normal temperature than would be expected at that temperature if the methane were an ideal gas. This effect is more marked at higher pressures. Similarly, as the pressure is increased at a constant temperature, more methane (measured at standard volume) can be stored in a given volume than would be predicted from the Ideal Gas Equation. "Standard volume" is volume measured at standard pressure and temperature (STP)).

Natural gas, like methane, shows z factor changes with pressure. Under about 1000 psia the dominant variable in the power relationship is the molecular weight of the gas. At this pressure level, addition of further amounts of ethane or propane increases the molecular weight of the gas more rapidly than the z factor decreases. Thus, there is an advantage to removing ethane and propane from the gas.

It is usual in the gas transportation and storage industry to try to strip out higher hydrocarbons such as ethane, propane, butane and unsaturated hydrocarbons from natural gas if the gas is to be transmitted through pipelines. This leaves mostly methane (with some traces of nitrogen and carbon dioxide) to be transported by the gas

pipeline. The materials which are stripped out are then transported or stored separately, often as liquids.

Summary of the Invention

It has now been found that, at pressures over 1,000 psia, it is advantageous to add to natural gas an additive which is a C_2 or C_3 hydrocarbon compound or a mixture of such additives. Above a lower limit (which varies with the additive being added and the pressure), this results in a smaller product of the z factor times the average molecular weight of the gas (hereinafter called the zM_w product) than would exist with methane alone, therefore leading to a decrease in the amount of power needed to pump the mixture or to compress it.

Detailed Description of the Invention

If ethane is the additive, enough ethane must be added to methane or natural gas to give a gas composition having a minimum of about 26% ethane for operation at 1,000 psia and normal temperatures (-40°F to +110°F). (All percentages in this document are percentages by volume). Ethane can be added until just before the mixture separates into separate gas and liquid (which occurs at about 40% ethane for a pressure 1,000 psia and a temperature of about 35°F). To reduce the danger of liquefaction if there is inadvertent pressure drop, and to reduce temperature extremes, generally operation at 26-35% ethane and 35°F to +40°F is preferred.

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When the pressure is raised to 2,200 psia, the addition of enough ethane to natural gas to give a gas composition having more than 6% ethane gives some beneficial results. Thus, as pressure increases, in the range from 1,000 psia to about 2,200 psia the beneficial results occur with less and less ethane. For the most beneficial results, however, an addition of enough ethane to give at least 15% ethane is preferred at pressures of 2,200 psia.

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Thus, for ethane as an additive, an amount is added to give a gas which has at least 26% ethane (but preferably 35% ethane) at 1,000 psia, and at least 6% ethane (but preferably 15% ethane) at 2,200 psia, with the minimum percentage of ethane decreasing smoothly with rise in pressure. Ethylene may be substituted for all

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or part of the ethane on a 1:1 volume basis. Where pressure fluctuates, as in a gas pipeline where the gas is compressed at compressor stations and becomes less compressed as it flows between compressor stations, the pressure indicated is the maximum pressure to which the gas is compressed. In such a compression-rarefaction arrangement, it is preferred that the ratio between the most compressed and the most rarefied pressures of the gas not exceed 1.3:1.

 C_3 hydrocarbons alone can also be used as the additive. Minimum useable percentage of the total gas mixture vary from a minimum of 5% at 1,000 psia to about 3% at 2,200 psia. Maximum amounts are those which will not cause separation of a liquid phase at the temperature used. The C_3 hydrocarbons may be any of propane, isopropane or propylene, separately or in admixture.

One or more C_3 hydrocarbons may also be substituted, preferably on a 1:3.5 volume basis, for C_2 hydrocarbons, but not to a point where they cause separation of a liquid phase at the pressure and temperature of operation. (A 1:3.5 basis means that each standard volume of C_3 hydrocarbon replaces three and a half standard volumes of C_2 hydrocarbon.) Generally, the limitation that a liquid phase should not be formed means that not more than about 12% of C_3 hydrocarbons should be present at 1000 psia and 60°F, and lesser amounts should be used as the pressure or temperature increases.

Two or more of the C_2 or C_3 additives can also be used. The use of two or more additives has a synergistic effect in many cases, so that less than the minimum amount of each is needed than would be needed if only one were present, in order to produce a zM_w product smaller than that of an equivalent standard volume of natural gas at the pressure and temperature involved.

At pressures over 1000 psig, C_4 hydrocarbons do not contribute much to the improvement of the zM_w product. Thus C_4 hydrocarbons are not additives contemplated by this invention. However C_4 hydrocarbons which are already present in the natural gas need not be removed if they are present in insufficient quantity to liquefy or to affect the zM_w product very adversely. The presence of more than 1% C_4 hydrocarbon in the mixture is not preferred, however, as C_4 hydrocarbons tend to liquify easily at pressures between 1,000 psia and 2,200 psia, and more than 1% C_4

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hydrocarbons give rise to increased danger that a liquid phase will separate out. C_4 hydrocarbons also have an unfavourable effect on the mixture's z factor at pressures just under 900 psia, so care should be taken that, during transport through a pipeline, that mixtures according to the invention which contain C_4 hydrocarbons are not allowed to decompress to less than 900 psia, and preferably not to less than 1,000 psia.

The addition of amounts of additive below the lower limit (unless two additives with a synergistic effect are used) actually increase the zMw product over that of methane or natural gas alone, and is thus detrimental. For example, when the pressure is 1,000 psia and the temperature is 35°F, mixtures of methane and ethane having less than about 26% ethane have a z factor greater than methane alone (all percentages are based on volumes at standard pressures and temperatures). Adding ethane to increase the percentage of ethane from 2% to, for example 12% at this pressure and temperature is therefore counterproductive, as it increases the zM, product and therefore increases the energy required to pump or compress for storage a given standard volume of gas. However, when more than 26% of ethane is present, the z factor falls so much that the zM, product tends to lower values than that of pure methane. The z factor continues to get smaller with increased percentages of ethane, bringing with it a lower zM, product to the point where further increase of ethane causes separation of a liquid phase (at about 40% ethane at 1,000 psia and 35°F). Thus, adding ethane to natural gas so that there is a mixture containing more than 26% ethane at 1,000 psia and 35°F leads to increased packing of molecules and a decreased zMw product, hence decreased pumping costs and more ability to store within a given volume. At 1,350 psia and 85°F, improved results over methane alone are obtained when only 17% of ethane is present in the mixture. Where the pressure is increased to 1,675 psia at 35°F, mixtures with 13% or more ethane, and the balance methane give improved results over methane alone. At 2,140 psia and 35°F, the improved effect is shown in mixtures of 6% ethane and the balance methane.

By the avoidance of liquid phase in this disclosure is meant the avoidance of enough liquid to provide a coherent liquid phase in the pipeline at the temperatures and pressures used. Such a phase can create pipeline problems through pooling in

low portions of the pipeline or forming liquid slugs which affect pumping efficiency. A few liquid droplets in the line however, can be tolerated.

The use of a hydrocarbon as additive has the additional advantage that it permits transport of a mixture of methane or natural gas and the hydrocarbon additive in a single pipeline with less energy expenditure than if the two were transported separately in separate pipelines.

Brief Description of the Drawings

The invention will be described further in association with the following drawings in which:

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Figures 1A to 1E are plots of capacity gain in percent against the content of C₂ hydrocarbons in a mixture of methane and ethane. Each of the plots shows the results at a different pressure. Figure 1F shows calculations of flow ratios for various mixtures of methane and ethane..

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Figures 2A and 2B are plots of capacity gain versus temperature (in degrees Fahrenheit) for pipelines at 800 psia and 1,675 psia respectively.

Figure 3 is a summary of pipeline horsepower requirements for various gas mixes, using a 36" pipeline operating at a maximum operation pressure of 1,740 psia, an inlet temperature of 80°F and a ground temperature of 32°F.

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Figure 4 is a plot of the horsepower requirements per thousand cubic feet of gas showing different mixtures of ethane and methane, at different pressures.

Detailed Description of the Embodiments Shown in the Drawings

Dealing first with Figure 1, this shows, for various pressures and the same temperature, the effect of the addition of ethane to methane. In each case, the z factor has been calculated for each percentage of ethane from 0 to 40%. Then, the lowest calculated z factor has been arbitrarily defined as 0% capacity gain. Each of the other results has been plotted as a percentage capacity gain with reference to the 0% capacity gain in order to prepare a curve. Curves developed in this way are given in Figures 1A to 1E, for different pressures, with each curve representing the situation for one pressure. The temperature represented by each curve is 35°F.

Looking at Figure 1A, it is seen that, for an 800 psia pipeline, the best packing occurs when the line is filled with pure methane. As ethane is added, the capacity gain percent decreases until there is about 25% ethane in the line. After this, the capacity gain begins to increase again, but it does not reach the levels obtained for pure methane.

Figure 1B shows the effect of addition to methane of ethane for a pipeline at 1,150 psia. Here, the capacity gain steadily decreases from 0% ethane to about 12% ethane, and then increases again. After approximately 25% ethane, the capacity gain is greater than occurred with no ethane at all.

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Figure 1C shows that this effect is even more pronounced when the pipeline pressure is increased to 1,350 psia. The lowest capacity now occurs at approximately 7%, and mixtures with more than 17% ethane exhibit packing (and hence, pipeline or storage capacity) gains unattainable with of natural gas or methane alone.

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Figure 1D shows that at 1,760 psia, the lowest capacity occurs at about 5%, and anything over 12% ethane gives a better capacity gain than is attainable with natural gas or methane alone. For best results, however, at least 15% ethane should be present.

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At 2,140 psia (Figure 1E) the addition of about 4% ethane gives a benefit, and the benefit steadily increases all the way up to the point at which the ethane begins to separate out in a liquid phase. For best results, however, at least 12% ethane, and preferably 15% ethane, should be present.

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Figure 1F shows the effect of the z factor and its product with the average molecular weight of the gas for gases having various amounts of methane and ethane (the ethane is shown in the table as " C_2 ") The flow ratio (1/the root of the zM $_{\rm w}$ product) is plotted.

Thus, it will be seen that for pressures above about 1,000 psia better packing, and hence lower compression cost and pumping cost for transportation occurs when increased amounts of ethane are added over a minimum amount which decreases with increasing pressure. At 1,150 psia (Figure 1B) about 24% ethane must

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be added to get the same packing effect as the approximately 2% ethane in normal natural gas. If more than this amount is added, however, better packing occurs with each addition. As the pressure increases, successively less ethane need be added to give better packing (and hence lower transmission costs and compression costs) than with natural gas. Indeed, at 2,410 psia, even about 4% ethane shows some advantage, although the advantage is of course greater as more ethane is added.

Figure 2 shows how the effect changes with temperature. Even at 800 psia (Figure 2A) there is a capacity increase (i.e. better packing) as temperature drops, and the capacity gain is greater the more C₂ that is present. However, the effect is not nearly as significant as at higher pressures. With higher pressure (Figure 2B) the capacity gain is much greater with temperature, and the improvement in capacity gain becomes still greater as increased amounts of ethane are added. Generally, therefore, it is preferred to operate at a relatively low temperature, such as 70° to -20°F. Higher temperatures (e.g. up to about 120°F) can be tolerated, but detract from the benefits of the invention.

Figure 3 shows horsepower required for different gas mixes of ethane and methane, through a pipeline at a maximum pressure of 1,740 psia and a minimum pressure of 1,350 psia. Figures are for a pipeline of 36" in diameter and a length of 1,785 miles, with pumping stations located every 56 miles. At a throughput of 2.0 million, standard cubic feet per day, a mixture of 98% methane and 2% ethane (which corresponds to ordinary natural gas) would require 812,579 horsepower. However, the same standard volume of gas, but containing 35% C₂, can be moved with only 661,860 horsepower, for a saving of over 150,000 horsepower. When the throughput is raised to 2.5 million standard cubic feet, natural gas containing 2% ethane cannot practically be transmitted, because the velocities and temperatures involved are too great. However, gas with 6% ethane can be transmitted, and gas with 35% ethane shows a saving of over 500,000 horsepower over that which is used to transmit the gas with 6% ethane.

Figure 4 shows the effect on horsepower requirements per million cubic feet of gas being pumped through the same pipeline as used in Figure 3 when the pipeline gas contains different concentrations of ethane at 35°F.

Figure 4 also shows the negative effect of adding ethane to a typical pipeline running at about 800 psia pressure and 35°F. Required power for pumping increases until the mix contains 26% ethane and then decreases for higher concentrations approaching the liquid phase limits. However, the decrease is not sufficient so that, by the concentration where liquefaction occurs (about 40%) there is any saving of horsepower over pumping ordinary natural gas. This energy hill however peaks at decreasing concentrations of ethane as operational pressure increases, e.g., 14% at 1,150 psia, 8% at 1,350 psia, 6% at 1,475 psia. This is due to the rate of decrease in the value of the z factor overcoming the rate of increase in density.

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As noted, at 800 psia, increasing the amount of ethane even up to 40% of the mixture does not produce a saving of horsepower. At 1,150 psia, however, an increase of ethane to over about 24% causes a horsepower saving when compared to the values at 2% ethane (which is approximately the amount of ethane in most natural gas). At 1,350 psia, a decrease in horsepower occurs with mixtures containing more than about 14% ethane. As the pressure increases, the horsepower saving occurs with still less added ethane. As can be seen from Figure 4, the peak horsepower requirement shifts to the left of the graph and becomes smaller as pressure increases. However, reading Figure 4 with Figure 2A and 2B shows that, as temperature decreases, the curve tends to rotate to the right. Thus, as temperature decreases, less addition of ethane is necessary to obtain the desired additional packing and hence decreased compression and pumping energy requirements. The result is lower compressive power requirements for equivalent volumes of higher B.T.U. gas mixes than can be achieved with conventional pipeline specification natural gas. Higher pipeline throughput are able to be economically achieved with these enhanced gas mixes with corresponding increases in horsepower.

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Similar effects are seen when ethylene is used in substitution for all or part of the ethane.

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When propane, isopropane or propylene, or mixtures of any of these gases, are substituted for all or part of the ethane, the effects are even more pronounced than with ethane, and occur with a smaller addition of each of the

compounds mentioned than was necessary with ethane as noted previously in this disclosure.

The preferred composition of the resulting gas is as follows:

COMPONENT	MAXIMUM VALUE	MINIMUM VALUE
METHANE	92% VOLUME	68% VOLUME
ETHANE AND/Or ETHYLENE	35% VOLUME	6% VOLUME
PROPANE AND/OR OTHER C ₃	12% VOLUME	0% VOLUME
BUTANES AND OTHER COMPONENTS OF THE NATURAL GAS	Not required, but amount present in original natural gas (up to about 1%)can be tolerated if it does not cause separation of a liquid phase at the pressure and temperature used	0% VOLUME
NITROGEN	Not required, but amount present in the original natural gas can be tolerated if below 2% by volume	0% VOLUME
CARBON DIOXIDE	Not required, but amount present in the original natural gas can be tolerated if below 1% by volume.	0% VOLUME
COMPONENTS TOTAL 100%		
TEMPERATURE	120°F	-20°F

20 Compositions with hydrocarbon additives outside these ranges generally are of little economic benefit, or approach the limits at which two-phase or liquid

1150 PSIA

2160 PSIA

PRESSURE

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behaviour can be seen. Therefore, if mixtures outside these parameters are used, care should be taken to avoid conditions which might cause liquid to fall out of the mixture. Liquid is generally to be avoided, as it may pool in low areas of a pipeline and be difficult to remove. The preferential liquefaction of some components will also cause the composition of the gaseous phase of the mixture to change, thereby changing the z factor and hence the compressibility of the gaseous phase.

The foregoing has illustrated certain specific embodiments of the invention, but other embodiments will of course be evident to those skilled in the art. Therefore it is intended that the scope of the invention not be limited by the embodiments described, but rather by the scope of the appended claims.

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What is claimed is:

- A method of transporting natural gas by pipeline, which comprises:
 - (a) adding to such natural gas sufficient of at least one C_2 or C_3 hydrocarbon or a mixture of C_2 and C_3 hydrocarbons such so the hydrocarbon, together with the C_2 and C_3 hydrocarbon (if any) originally in the natural gas, forms a resulting mixture with a total C_2 or C_3 hydrocarbon content which is sufficient, at the pressure and temperature to be used for transporting, to reduce the product of the z factor and the average molecular weight of the resulting mixture to a level lower than that of the untreated natural gas, and
- (b) transporting such resulting mixture by pipeline at a temperature of between -40° and +120° Fahrenheit and pressure greater than 1000 psia, said pressure and temperature being chosen so the resulting mixture has no liquid phase at the temperature and pressure of transmission.
 - 2. A method as claimed in claim 1 where the hydrocarbon is selected from
- (a) between 26 and 40% of at least one C₂ compound if the pressure is about 1000 psia, declining smoothly to about 6% to 15% of said C₂ compound if the pressure is about 2200 psia, or
 - (b) between 12% and 4% of a C_3 compound, if the pressure is about 1000 psia, declining smoothly to the C_3 amount which will not cause liquefaction at the pressure used when the pressure is above 1000 psia.
 - 3. A method as claimed in either claim 1 or claim 2, in which there is not more than 1% by volume of carbon dioxide in the resulting mixture.

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- 4. A method as claimed in claim 1 or claim 2, in which there is not more than 2% nitrogen in the resulting mixture.
- 5. A method as claimed in any of claims 1-4, in which the temperature at which the resulting mixture is transmitted is between -20° F. and +120°F.
- 5 6. A method as claimed in any of claims 1-4, in which the pressure at which the resulting mixture is transmitted is between 2160 psia and 1150 psia.
 - 7. A method as claimed in any of claims 1-6 in which the C_2 hydrocarbon added to the natural gas is ethane.
- 8. A method as claimed in any of claims 1-7, in which the C₃ hydrocarbon added to the natural gas is propane.
 - 9. A gas mixture, for use in a pipeline at a pressure greater than 1,000 psia and a temperature of from -40 degrees F. to +120 degrees F., which comprises:
 - (a) from 65 to 92% by volume of methane
 - (b) from 6 to 35% by volume of ethane
 - (c) from 0 to 9% by volume of propane
 - (d) from 0% by volume of C₄ hydrocarbons to a percentage of Cu hydrocarbons which does not liquefy at the pressure used
 - (e) not more than 1% each of carbon dioxide or nitrogen the total being 100%, and such mixture being completely gaseous with no liquid phase at the temperature and pressure of intended operation.

10. A gas mixture as claimed in claim 9, said gas mixture being at a pressure of 1000-2200 psia and a temperature of from -20 degrees F. to +120 degrees F.

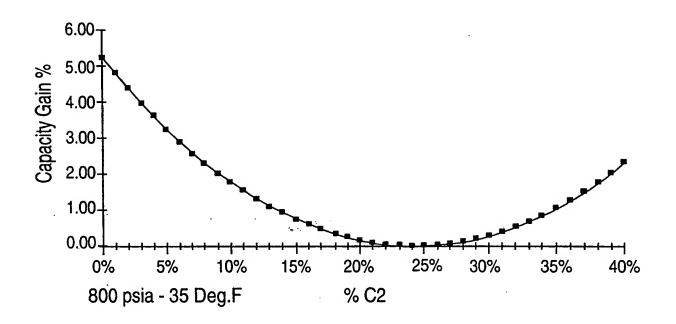


FIG. 1A

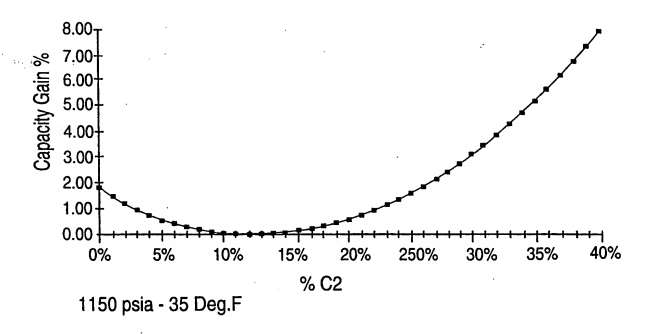


FIG. 1B

SUBSTITUTE SHEET (RULE 26)

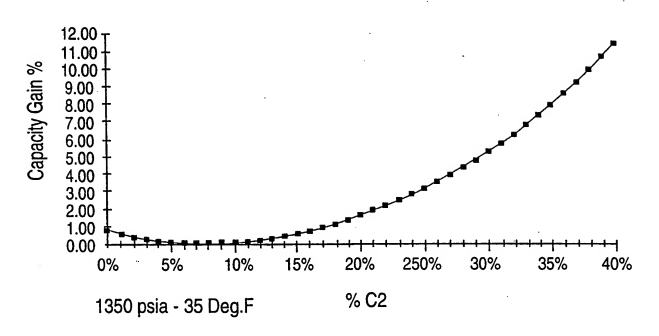


FIG. 1C

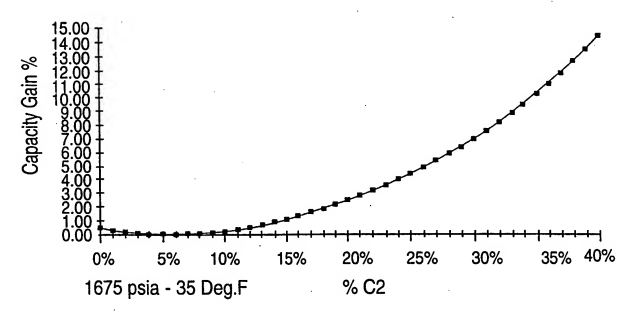


FIG. 1D

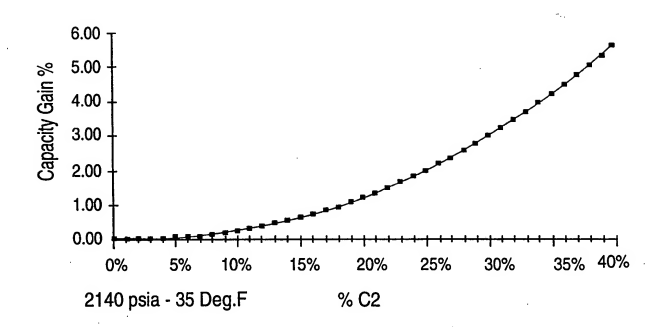


FIG. 1E

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TABLE A:		Fl	LOW RATIO	OS FROM	Mw & Z VALUES	*
					$\frac{1 @ 30^{0}F}{\sqrt{\text{(Mw.Z)}}}$	$\frac{1 @ 60^{0}F}{\sqrt{\text{(Mw.Z)}}}$
		Mw	Z@ 30 ⁰ F	Z@ 60 ⁰ F	(% Change from Base Case)	(% Change from Base Case)
Fig.A: Gas 100% C1 BaseCases	at 1100 psia	16.04	0.817	0.856	0.2763 (0.00%)	0.26699 (0.00%)
Fig.B: Gas 90% C1 10% C2	at 1100 psia	17.45	0.769	0.819	0.2730 (-1.19%)	0.2645 (-2.00%)
Fig.C: Gas 80% C1 20% C2	at 1100 psia	18.85	0.711	0.775	0.2732 (-1.12%)	0.2616 (-3.08%)
Fig.D: Gas 70% C1 30% C2	at 1100 psia	20.25	0.638	0.722	0.2782 (+0.69%)	0.2615 (-3.30%)

	-				$\frac{1 @ 30^{0}F}{\sqrt{\text{(Mw.Z)}}}$	$\frac{1 @ 60^{\circ}F}{\sqrt{(Mw.Z)}}$
		Mw	Z@ 30 ⁰ F	Z@ 60 ⁰ F	(% Change from Base Case)	(% Change from Base Case)
Fig.A: Gas 100% C1	at 1900 psia	16.04	0.742	0.798	0.2899 (+4.92%)	0.2795 (+3.37%)
Fig.B: Gas 90% C1 10% C2	at 1900 psia	17.45	0.681	0.746	0.2901 (+4.99%)	0.2772 (+2.70%)
Fig.C: Gas 80% C1 20% C2	at 1900 psia	18.85	0.616	0.689	0.2935 (+6.23%)	0.2775 (+2.82%)
Fig.D: Gas 70% C1 30% C2	at 1900 psia	20.25	0.553	0.628	0.2988 (+8.14%)	0.2804 (+3.89%)

FIG. 1F SUBSTITUTE SHEET (RULE 26)

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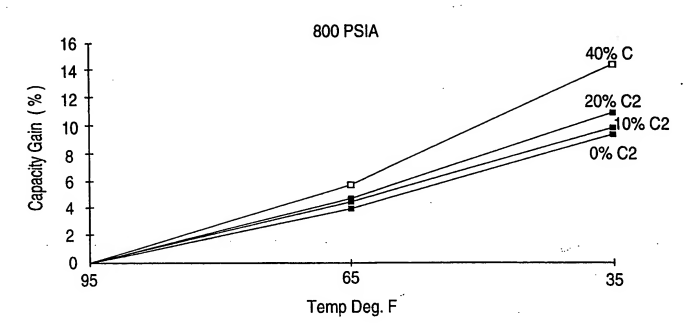


FIG. 2A

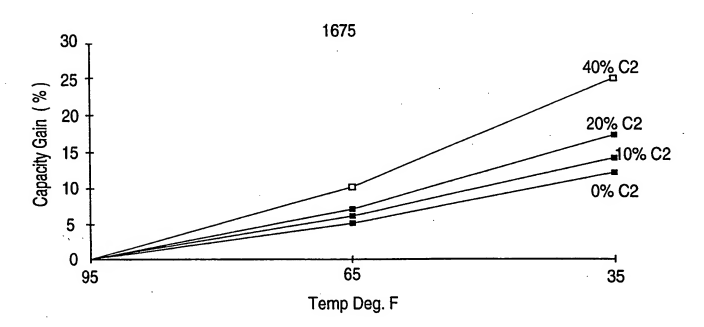


FIG. 2B

SUMMARY OF PIPELINE HORSEPOWER **REQUIREMENTS FOR GAS MIXES**

PIPELINE DIAMETER:

36"

INLET TEMPERATURE:

MAX.OPERATION PRESSURE: 1740 psia

GROUND TEMPERATURE: 32° F

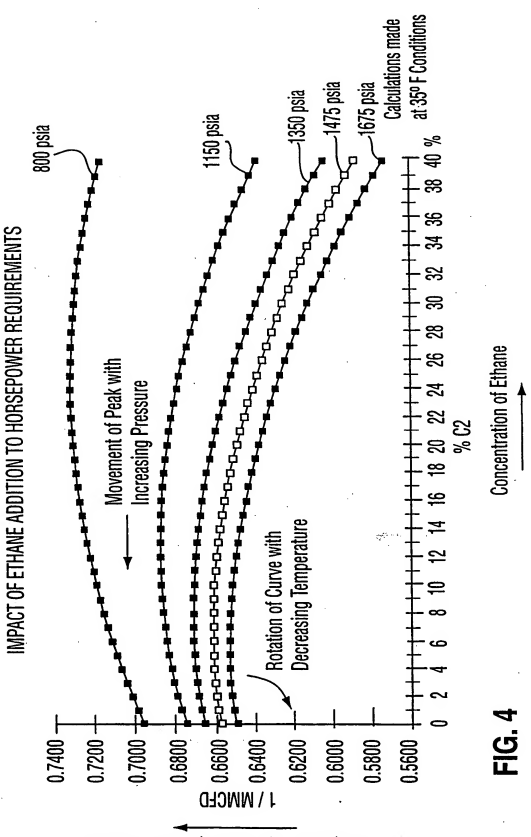
70.

THRUPUT MMSCF/D	TYPICAL TRANSMISSION GAS 2 % C ₂	ENHANCED GAS 6 % C ₂	ENHANCED GAS 14 % C ₂	ENHANCED GAS 35 % C ₂
500	10 549	10 235	9857	7724
1000	92 018	89 705	87 157	75 071
1500	343 621	330 406	315 833	271 784
2000	812 579	760 244	714 199	651 860
2300	1 480 446 *	* *	* *	* *
2500	. **	1 784 791	1 387 893	1 209 985

APPROACHING PRACTICAL LIMITS

FIG. 3

^{**} BEYOND PRACTICAL LIMITS



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INTERNATIONAL SEARCH REPORT

in atlonal Application No PCT/CA 98/00354

A. CLASSI IPC 6	FICATION OF SUBJECT MATTER C10L3/06 F17D1/00 F17D1/04	F17D1/17 F17D1	/02
	o International Patent Classification (IPC) or to both national classificat	tion and IPC	
	SEARCHED currentation searched (classification system followed by classification	n symbola)	
IPC 6	C10L F17D		
Documenta	tion searched other than minimum documentation to the extent that su	ch documents are included in the fields sea	rched
Ejectronic	ata base consulted during the international search (name of data bas	e and, where practical, search terms used)	
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the rele	vant passages	Relevant to claim No.
P,X	WO 97 19151 A (665976 ALBERTA LTD IAN (CA); PERRY GLEN (CA)) 29 May see claims 1-10		1-10
Α .	US 5 315 054 A (TEEL DALE) 24 May	1994	
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Furt	her documents are listed in the continuation of box C.	X Patent family members are listed in	n annex.
* Special ca	tlegories of cited documents:	"T" later document published after the inter	national filing date
consi	ent defining the general state of the art which is not dered to be of particular relevance	or priority date and not in conflict with cited to understand the principle or the invention	the application but
filing of the country	ant which may throw doubts on priority claim(s) or	"X" document of particular relevance; the c cannot be considered novel or cannot involve an inventive step when the do	be considered to
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"P" docum	means ent published prior to the international filing date but han the priority date claimed	ments, such combination being obviou in the art. "&" document member of the same patent	
Date of the	actual completion of the international search	Date of mailing of the international sear	rch report
3	1 August 1998	07/09/1998	
Name and	mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk	Authorized officer	
	Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	De Herdt, O	

INTERNATIONAL SEARCH REPORT

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